WEST Search History

Hide Items Clear Cancel Restore

DATE: Wednesday, June 22, 2005

Hide?	<u>Set</u> Name	Query								
	DB=PC	GPB, USPT, EPAB, JPAB; PLUR=YES; OP=OR								
T	L22	(acyl adj transferase) same gene	169							
	L21	L20 and (transacylase or acyltransferase or transferase)								
	L20	1999	60							
	L19	L18 same (transform\$6 or transgen\$4 or exogenous)	398							
	L18	triacylglycerol or triglyceride) same (produc\$6 or make or \$4synthesi\$5)								
<u>. </u>	L17	revisiae same (triacylglycerol or triglyceride) same (produc\$6 or make or 4synthesi\$5)								
	L16	cerevisiae same (triacylglycerol or triglyceride) same (transform\$6 or transgen\$4 or exogenous)	5							
	L15	1999	6							
	L14	L13 same(express\$4 or transcri\$5 or \$6produc\$4)	138							
	L13	L12 same (DGAT or ACAT or diacyl\$10)	182							
. С	L12	(transacylase or acyltransferase or transferase)same (triacylglycerol or triglyceride)	636							
	L11	L10 and triacylglycerol	576							
	L10	(transacylase or acyltransferase or transferase)	41496							
	L9	1999	7							
	L8	1999	4							
	L7	L6 same (express\$4 or transcri\$5 or \$6produc\$4)	74							
	L6	L4 same (DGAT or ACAT or diacyl\$10)	107							
To appropriate to	L5	L4 same (DGAT or ACAT)	25							
	L4	(transacylase or acyltransferase or transferase) same triacylglycerol	188							

END OF SEARCH HISTORY

STN SEARCH SUMMARY 09/937779

=> d his

	FILE	'CAPL	JS	' EN	NTER	ED AT 14:36:18 ON 22 JUN 2005
L2		58760	S	(TF	RANS	ACYLASE OR ACYLTRANSFERASE OR TRANSFERASE)
L3		11564	S	TRI	[ACYI	LGLYCEROL
L4		770	S	L2	AND	L3
L5		508	S	L4	AND	PD<1999
L6		1451	S	L5	AND	DAGAT OR ACAT
L7		['] 39	S	L5	AND	(DAGAT OR ACAT)
L8		574	S	L4	AND	(DAGAT OR ACAT OR DIACYLGLYCEROL OR CHOLESTEROL)
L9		24	S	L8	AND	CEREVISIAE

SEQUENCE STARCH SUMMARY 09/937779

GenCore version 5.1.6 Copyright (c) 1993 - 2005 Compugen Ltd.

OM nucleic - nucleic search, using sw model

Run on:

May 2, 2005, 13:58:51; Search time 8752 Seconds

(without alignments)

10995.436 Million cell updates/sec

Title:

US-09-937-779-1

Perfect score:

Sequence:

1 atgggcacactgtttcgaag.....agatgcccttcccaatgtaa 1986

Scoring table:

IDENTITY NUC

Gapop 10.0 , Gapext 1.0

Searched:

4708233 segs, 24227607955 residues

Total number of hits satisfying chosen parameters:

9416466

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database :

GenEmbl:*

1: gb ba:*

gb htg:*

gb in:*

4: gb om:*

gb_ov:* 5:

gb pat:*

7: gb_ph:*

8: gb pl:*

gb_pr:*

10: gb ro:*

11: gb sts:*

12: gb_sy:*

13: gb_un:*

14: gb_vi:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

Result No.			Length				Description
1	1986	100.0	1986	6	BD271616 BD271625	2000	BD271616 Novel cla
2	1986	100.0	1986	6	BD271625	Appl	BD271625 Novel cla
3	1986	100.0	1986	6	BD271626	17	BD271626 Novel cla

```
AX037577 WU 2000
                100.0
                        1986
         1986
                                                              AX037577 Sequence
                                  AX037595 //
AX037597 //
    5
         1986
                100.0
                        1986
                               6
                                                              AX037595 Sequence
    6
         1986
                100.0
                        1986
                               6
                                  AX090382 WO 200 1
                                                              AX037597 Sequence
                                  SCYNROOSW 4/1996 Stepped AX090382 Sequence
    7
         1986
                100.0
                        1986
                               6
                                  SCN201952 1994 Verhasselt
    8
         1986
                100.0
                        2575
                                                              Z71623 S.cerevisia
    9
         1986
                100.0 23901
                                                              X77395 S.cerevisia
                                  CR380955 05 2004
   10
        835.6
                 42.1 110000
С
                               8
                                                              Continuation (6 of
   11
        715.2
                 36.0 110000
                                  CR382126 06
С
                                                              Continuation (7 of
С
  12
        676.2
                 34.0 110000
                               8
                                  AE016819 00
                                                              AE016819 Eremothec
  13
        676.2
                 34.0 110000
                               8
                                  AE016819 01
                                                              Continuation (2 of
   14
        555.6
                 28.0 110000
                                  CR382136 07
                                                              Continuation (8 of
   15
        555.6
                 28.0 110000
                               8
                                  CR382136 08
                                                              Continuation (9 of
          537
                        2106
   16
                 27.0
                               6
                                  AR546016
                                                              AR546016 Sequence
  17
        439.8
                 22.1 110000
                                  CR382131 19
                                                              Continuation (20 o
                               8
   18
        273.4
                 13.8 103568
                                  CNS07EGJ
                                                              AL590462 DNA centr
  19
        273.4
                 13.8 322194
                               8
                                  CNS09S4S
                                                              BX088700 DNA centr
  . 20
          202
                 10.2
                        2312
                               6
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                                                              BD271617 Novel cla
   21
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                        2312
                               6
                                  BD271627
                                                              BD271627 Novel cla
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                                                              AX037599 Sequence
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                               8
                                  SPBC776
                                                              AL035263 S.pombe c
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        137.2
                  6.9
                        2441
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                                  BT013705
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                                  AX794706
                                                              AX794706 Sequence
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                  5.9
                         2013
                               6
                                  AX090380
                                                              AX090380 Sequence
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                  5.9
                        2016
                               6
                                  AX412864
                                                              AX412864 Sequence
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                               6
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                                                              AX794714 Sequence
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                                                              AX794710 Sequence
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                                  AX794704
                                                              AX794704 Sequence
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                                  AX794722
                                                              AX794722 Sequence
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                  5.1
                        1945
                               8
                                  PPA493274
                                                              AJ493274 Physcomit
   44
         82.4
                  4.1
                       87724
                               8
C
                                  AY448010S4
                                                              AY448013 Ipomoea t
   45
         67.6
                  3.4
                        2914
                               8
                                  PPA493276
                                                              AJ493276 Physcomit
```

OM nucleic - nucleic search, using sw model

Run on: May 2, 2005, 11:44:51; Search time 1103 Seconds

(without alignments)

10658.755 Million cell updates/sec

Title: US-09-937-779-1

Perfect score: 1986

Sequence: 1 atgggcacactgtttcgaag.....agatgcccttcccaatgtaa 1986

Scoring table: IDENTITY NUC

Gapop 10.0 , Gapext 1.0

Searched: 4390206 seqs, 2959870667 residues

Total number of hits satisfying chosen parameters: 8780412

Minimum DB seg length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database: N Geneseq 16Dec04:*

1: geneseqn1980s:*

2: geneseqn1990s:*

3: geneseqn2000s:*

4: geneseqn2001as:*

5: geneseqn2001bs:*

6: geneseqn2002as:*

7: geneseqn2002bs:*

8: geneseqn2003as:*

9: geneseqn2003bs:*

10: geneseqn2003cs:*

11: genesegn2003ds:*

12: geneseqn2004as:*

13: geneseqn2004bs:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

Result No.	Score	Query Match	Length	DB	ID	Description
1	1986	100.0	1986	3	AAC64431 Appl	Aac64431 Saccharom
2	1986	100.0	1986		AAC64440 //	Aac64440 Saccharom
3	1986	100.0	1986	5	AAS01342 WO 0	Aas01342 Yeast LCA
4	1986	100.0	1986	12	ADF47816 April 2002	Adf47816 Yeast acy

						,	
	5	1958.8	98.6	1986	3	AAC64441 App	Aac64441 Saccharom
	6	1694.4	85.3	1701	12	ADF47817	Adf47817 Yeast mem
	7	309.2	15.6	1872	12	ADF47831	Adf47831 Schizosac
	8	254	12.8	2223	12	ADF47830	Adf47830 Aspergill
С	9	238	12.0	7299	13	ADR84383 2003	Adr84383 Aspergill
-	10	217.6	11.0	2047	12	ADF47829/4/p	Adf47829 Aspergill
	11	202	10.2	2312	3	AAC64442 //	Aac64442 Schizosac
	12	202	10.2	2312	3	AAC64432 //	Aac64432 Schizosac
	13	133.8	6.7	2004	12	ADF47835-/	Adf47835 Crepis pa
	14	127.2	6.4	2433	9	AAL62907 2001	Aal62907 Rice cDNA
	15	126.6	6.4	2016	12	ADF47833 2002	Adf47833 Crepis pa
	16	124	6.2	1998	8	ABZ76360	Abz76360 A. thalia
	17	124	6.2	1998	12	ADF47827	Adf47827 Arabidops
	18	124	6.2	2398	9	AAL62909	Aal62909 Sunflower
	19	117.6	5.9	2565	9	AAL62906	Aal62906 Corn cDNA
	20	117.4	5.9	1803	12	ADF47821	Adf47821 Arabidops
	21	117.4	5.9	2013	5	AAS01341	Aas01341 Arabidops
	22	117.4	5.9	2016	6	ADG88186	Adg88186 A. thalia
	23	117.4	5.9	2016	12	ADF47819	Adf47819 Arabidops
	24	117.4	5.9	2425	10	ADG25155	Adg25155 P. patens
	25	117.4	5.9	2427	3	AAC64434	Aac64434 Arabidops
	26	117	5.9	553	8	ABZ54921	Abz54921 Aspergill
	27	116.6	5.9	2030	9	AAL62910	Aal62910 Wheat cDN
	28	115.4	5.8	2700	9	AAL62908	Aal62908 Soybean c
	29	114.4	5.8	2479	9	AAL62905	Aal62905 Guayule c
	30	105.4	5.3	4093	9	AAL62913	Aal62913 Soybean c
	31	103.4	5.2	578	13	ADR61378	Adr61378 Cotton cD
	32	63.8	3.2	419	5	AAS01089	Aas01089 Soybean s
	33	63.4	3.2	3107	5	AAS01085	Aas01085 Arabidops
	34	63.4	3.2	3685	3	AAC64433	Aac64433 Arabidops
	35	63.4	3.2	3685	3	AAC64443	Aac64443 Arabidops
	36	59.2	3.0	616	3	AAC64436	Aac64436 Neurospor
	37	58.6	3.0	254	5	AAS01104	Aas01104 Corn ster
	38	58.4	2.9	1510	9	AAL62904	Aal62904 Guar cDNA
С	39	54.8	2.8	1680	5	AAS01086	Aas01086 Arabidops
С	40	51	2.6	2000	8	ADA71938	Ada71938 Rice gene
	41	49.4	2.5	1641	5	AAS01082	Aas01082 Arabidops
	42	49.4	2.5	1902	12	ADF47823	Adf47823 Arabidops
	43	45.6	2.3	2000	8.	ADA71938	Ada71938 Rice gene
С	44	44.2	2.2	4590	5	AAH24065	Aah24065 Yeast AOD
	45	44	2.2	655	13	ADR63567	Adr63567 Cotton cD

OM nucleic - nucleic search, using sw model

Run on: May 2, 2005, 15:50:07; Search time 355 Seconds

(without alignments)

9153.931 Million cell updates/sec

Title: US-09-937-779-1

Perfect score: 1986

Sequence: 1 atgggcacactgtttcgaag.....agatgcccttcccaatgtaa 1986

Scoring table: IDENTITY NUC

Gapop 10.0 , Gapext 1.0

Searched: 1202784 seqs, 818138359 residues

Total number of hits satisfying chosen parameters: 2405568

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database : Issued_Patents NA:*

1: /cgn2 6/ptodata/1/ina/5A COMB.seg:*

2: /cgn2 6/ptodata/1/ina/5B COMB.seq:*

3: /cgn2 6/ptodata/1/ina/6A COMB.seq:*

4: /cgn2 6/ptodata/1/ina/6B COMB.seq:*

5: /cgn2 6/ptodata/1/ina/PCTUS COMB.seq:*

6: /cgn2 6/ptodata/1/ina/backfiles1.seq:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

			*			•		
Res	ult		Query					
	No.	Score	Match	Length	DB	ID	Description	on 4747-137
	1	537	27.0	2106	4	US-09-248-796A-1147 FD=2/9 US-08-232-463-14 X US-09-806-708B-22	9 Sequence	1147, Ap Patent 102
С	2	. 55	2.8	7218	1	US-08-232-463-14 🔏	Sequence	14, Appl
	3	47.4	2.4	1141	4	US-09-806-708B-22⊁	Sequence	22, Appl /
С	4	42.2	2.1	1141	4	US-09-806-708B-22 X	Sequence	22, Appl /
	5	38.6	1.9	1431	4	US-09-248-796A-9904 PD 72/9	/ Sequence	9904, Ap
С	6	38.6	1.9	24333	4	US-09-639-207-9 √	Sequence	9, Appli /
	7	38	1.9	601	4	US-09-949-016-149237X	Sequence	149237,
С	8	38	1.9	187169	4	US-09-949-016-12776 [.] X		12776, A
С	9	38	1.9	191569	4	US-09-949-016-15940 🗸 🚄		15940, A
	10	37.4	1.9	552	4	US-09-248-796A-103772/97		10377, A
	11	37.2	1.9	615	4	us-09-134-000c-1430 1998	Sequence	1430, Apbadend
						_		10710

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Sequence 4906, Ap 6747137
                                  US-09-248-796A-4906 2/90
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                                  US-09-949-016-14319 X
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                                                                Sequence 14319, A
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                         8905
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                                                                Sequence 11761, A
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                                  US-09-949-016-16261/
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                                                                Sequence 16261, A
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                                  US-09-949-016-12902
                                                                Sequence 12902, A
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                  1.9 640681
                                  US-09-790-988-1
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OM nucleic - nucleic search, using sw model

Run on: May 2, 2005, 15:32:31; Search time 1201 Seconds

(without alignments)

10066.732 Million cell updates/sec

Title: US-09-937-779-1

Perfect score: 1986

Sequence: 1 atgggcacactgtttcgaag.....agatgcccttcccaatgtaa 1986

IDENTITY NUC Scoring table:

Gapop 10.0 , Gapext 1.0

Searched: 5642217 seqs, 3043843248 residues

Total number of hits satisfying chosen parameters: 11284434

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database : Published Applications NA:*

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2: /cgn2_6/ptodata/1/pubpna/PCT NEW PUB.seq:*

3: /cgn2 6/ptodata/1/pubpna/US06 NEW PUB.seg:*

4: /cgn2 6/ptodata/1/pubpna/US06 PUBCOMB.seg:*

/cgn2_6/ptodata/1/pubpna/US07_NEW PUB.seq:* 5:

/cgn2 6/ptodata/1/pubpna/PCTUS PUBCOMB.seg:*

/cgn2 6/ptodata/1/pubpna/US08 NEW PUB.seq:* 7:

/cgn2 6/ptodata/1/pubpna/US08 PUBCOMB.seq:*

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10: /cgn2 6/ptodata/1/pubpna/US09B PUBCOMB.seq:*

11: /cgn2 6/ptodata/1/pubpna/US09C PUBCOMB.seg:*

12: /cgn2 6/ptodata/1/pubpna/US09 NEW PUB.seq:*

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14: /cgn2_6/ptodata/1/pubpna/US10B PUBCOMB.seq:*

/cgn2_6/ptodata/1/pubpna/US10C_PUBCOMB.seq:* 15:

/cgn2_6/ptodata/1/pubpna/US10D PUBCOMB.seq:*

17: /cgn2 6/ptodata/1/pubpna/US10E PUBCOMB.seq:*

/cgn2 6/ptodata/1/pubpna/US10F PUBCOMB.seq:*

/cgn2_6/ptodata/1/pubpna/US10 NEW PUB.seg:*

/cgn2_6/ptodata/1/pubpna/US11_NEW_PUB.seq:* 20:

/cgn2_6/ptodata/1/pubpna/US60_NEW_PUB.seq:* 21:

/cgn2_6/ptodata/1/pubpna/US60 PUBCOMB.seg:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

Result Query No. Score Match Length DB ID Description ------------US-10-883-760-45 X 1 436.6 22.0 2326 19 Sequence 45, Appl 2 128.6 6.5 2395 18 US-10-437-963-38278 С Sequence 38278, A 3 127.2 6.4 2433 17 US-10-321-802-21X Sequence 21, Appl 4 124 6.2 1998 16 Sequence 5, Appli 5 124 6.2 2398 17 US-10-321-802-25X Sequence 25, Appl 18 6 117.8 5.9 2700 US-10-425-115-55049 X Sequence 55049, A 7 117.6 5.9 1388 18 С US-10-425-115-55048X Sequence 55048, A 8 117.6 5.9 2565 17 US-10-321-802-19X Sequence 19, Appl 9 116.6 5.9 2030 17 US-10-321-802-27× Sequence 27, Appl 10 115.4 5.8 17 2700 US-10-321-802-23X Sequence 23, Appl 11 114.4 5.8 2479 17 US-10-321-802-17X Sequence 17, Appl 12 105.4 5.3 4093 17 US-10-321-802-35 X Sequence 35, Appl 13 103.4 5.2 578 18 US-10-767-795-2159X Sequence 2159, Ap US-10-424-599-125713X 14 101.8 5.1 2838 17 Sequence 125713, 15 66.8 3.4 735 18 US-10-437-963-38267 Sequence 38267, A 16 58.4 2.9 1510 17 US-10-321-802-15 Sequence 15, Appl 17 56.2 2.8 1141 17 US-10-424-599-124418 Sequence 124418, 18 55.4 2.8 1218 17 US-10-425-114-18228 Sequence 18228, A 19 55.4 2.8 1239 18 US-10-425-115-71787 C Sequence 71787, A 20 53.2 2.7 887 18 US-10-767-701-9911 Sequence 9911, Ap 21 49 2.5 693 18 US-10-425-115-55043 Sequence 55043, A 22 44 2.2 655 18 US-10-767-795-4348 Sequence 4348, Ap 23 42 2.1 924 18 US-10-425-115-137717 Sequence 137717, 24 42 2.1 1383 18 US-10-425-115-93909 Sequence 93909, A 25 41.8 2.1 3673778 16 US-10-312-841-1 Sequence 1, Appli 26 41.6 2.1 1659 С 9 US-09-938-842A-4435 Sequence 4435, Ap 27 41.6 1659 2.1 11 US-09-938-842A-4435 Sequence 4435, Ap 28 41.6 2.1 2790 US-09-938-842A-698 Sequence 698, App 29 41.6 2.1 2790 11 US-09-938-842A-698 Sequence 698, App 30 41.4 2.1 468 18 US-10-674-124A-1427 Sequence 1427, Ap 31 2.1 41.4 599 17 US-10-338-110-117 Sequence 117, App 32 41.2 2.1 6271 15 US-10-172-086-36 Sequence 36, Appl 33 41.2 2.1 6271 15 US-10-311-455-1310 Sequence 1310, Ap 34 41.2 2.1 17 6271 US-10-221-714A-178 Sequence 178, App 35 41.2 2.1 6271 18 US-10-311-507-62 Sequence 62, Appl 36 41.2 2.1 6271 18 US-10-480-846-36 Sequence 36, Appl 37 41 2.1 848 18 US-10-425-115-93907 Sequence 93907, A 38 41 2.1 5314 15 US-10-311-455-133 Sequence 133, App 39 41 2.1 5314 16 US-10-240-452-9 Sequence 9, Appli 40 40.6 2.0 539 10 US-09-814-353-17862 Sequence 17862, A 41 40.6 2.0 6327 14 US-10-239-676-152 Sequence 152, App 42 40.6 2.0 6327 15 US-10-240-453-168 Sequence 168, App 43 40.4 2.0 540 17 US-10-424-599-5143 Sequence 5143, Ap 44 40 2.0 5879 15 US-10-311-455-242 Sequence 242, App 45 40 2.0 7201 15 US-10-311-455-310 Sequence 310, App

OM nucleic - nucleic search, using sw model

Run on: May 2, 2005, 15:27:01; Search time 6684 Seconds

(without alignments)

11309.940 Million cell updates/sec

Title: US-09-937-779-1

Perfect score: 1986

Sequence: 1 atgggcacactgtttcgaag.....agatgcccttcccaatgtaa 1986

Scoring table: IDENTITY NUC

Gapop 10.0 , Gapext 1.0

Searched: 34239544 seqs, 19032134700 residues

Total number of hits satisfying chosen parameters: 68479088

Minimum DB seg length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database : EST:*

1: gb_est1:*

2: gb_est2:*

3: gb htc:*

4: gb est3:*

5: gb est4:*

6: gb est5:*

7: gb est6:*

8: gb gss1:*

9: gb gss2:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

Res	ult No.	Score	% Query Match	Length	DB	ID	Description
	1	439	22.1	832	9	CNS060AQ(2000	AL407832 T7 end of
С	2	427.8	21.5	948	9	CNS06Y94 🛠	AL420734 T3 end of
	3	354.6	17.9	609	8	BZ297416X	BZ297416 CG3420.f1
С	4	324.8	16.4	400	8	AZ924376×	AZ924376 4906.ic28
	5	291.6	14.7	978	9	CNS06TH7X	AL414545 T7 end of
С	6	249.8	12.6	882	9	CNS07EA9	AL441511 T7 end of
С	7	241.2	12.1	474	8	AZ928292 🗙	AZ928292 479.dif06
С	8	235.4	11.9	509	8	BZ302540X	BZ302540 KD1890.q1

		9	208.2	10.5	783	7	C0026714 X	CO026714 EST805098
		10	199.6	10.1	512	8	BZ301369 X	BZ301369 KD1229.g1
	С	11	194.6	9.8	849		CNSO6V5M X	AL416720 T7 end of
	_	12	165.4	8.3	899		C0003381 X	CO003381 EST791716
		13	151.2	7.6	970		CO032392X	C0032392 EST810776
	С	14	134.2	8.3 7.6 6.8 6.5	550	7	CF884811 V	CF884811 tric042vm
	_	15	129.8	6.5	596	7	CN846346 X	CF884811 tric042xm CN846346 PG07016B0
		16	122.6	6.2	605		BE450991	BE450991 EST401878
		17	122.2	6.2	719	5	BQ865802	B0865802 0GC5021 v
		18	119.8	6.0	821	4	BM780050	BM780050 EST590626
		19	118.2	6.0	854	4		BG645669 EST507288
		20	117.4	5.9	867		CF821111	CF821111 EST698493
		21	116.6	5.9	1490		CA731511	CA731511 wiplc.pk0
		22	115.2	5.8	834		CF820302	CF820302 EST697684
		23	113.2	5.7	844		CB631429	CB631429 OSIIEb08P
		24	112.8	5.7	634		BQ870476	BO870476 OGD9C14.v
		25	108.8	5.5	586	1	AV940595	BQ870476 QGD9C14.y AV940595 AV940595
		26	108.8	5.5	610		BJ472152	B.1/1/2152 B.1/1/2152
		27	108.8	5.5	628	4	BJ478864	BJ478864 BJ478864
		28	108.8	5.5	674	4	BJ480170	BJ480170 BJ480170
		29	107.2	5.4	602	6	CA210413	CA210413 SCEPSB112
		30	107.2	5.4	724	5	BQ803421	BQ803421 WHE2837 D
,		31	106.2	5.3	569	6	CA018479	CA018479 HV08L24r
		32	104.2	5.2	566	4	BJ470786	BJ470786 BJ470786
		33	102.6	5.2	680	7	CN846719	CN846719 PG07019E0
		34	98.2	4.9	902	7	CF704247	CF704247 CCAC369TR
		35	95	4.8	575		AV938810	AV938810 AV938810
		36	94.6	4.8	568		CA184037	CA184037 SCQSST311
		37	89	4.5	578		BM062412	BM062412 KS01042D1
		38	87.6	4.4	448	6	CA018796	CA018796 HV09K23r
		39	85.8	4.3	631	2		BF053150 EST438380
		40	85.8	4.3	869	7		CF685137 CCAGP17TR
		41	85.6	4.3	626		AW587308	AW587308 EST318931
		42	85.6	4.3	1005		CD458189	CD458189 Fg08_10b0
		43	84.8	4.3	477		BG313245	BG313245 WHE2051_F
	С	44	83.6		885		CF820303	CF820303 EST697685
	C	45	83.4	4.2	943	7	CF821110	CF821110 EST698492

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OM nucleic - nucleic search, using sw model

Run on: May 2, 2005, 18:33:57; Search time 8752 Seconds

(without alignments)

10995.436 Million cell updates/sec

Title: US-09-937-779-1

Perfect score: 1986

Sequence: 1 atgggcacactgtttcgaag.....agatgcccttcccaatgtaa 1986

Scoring table: OLIGO NUC

Gapop 60.0 , Gapext 60.0

Searched: 4708233 seqs, 24227607955 residues

Word size : 0

Total number of hits satisfying chosen parameters:

9416466

Minimum DB seq length: 0

Maximum DB seq length: 200000000

Post-processing: Listing first 45 summaries

Database : GenEmbl:*

1: gb ba:*

2: gb_htg:*

3: gb_in:*

4: gb om:*

5: gb ov:*

6: gb_pat:*

7: gb ph:*

8: gb pl:*

9: gb pr:*

10: qb ro:*

11: gb sts:*

12: gb_sy:*

13: gb_un:*

14: gb_vi:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

Result		% Query				
No.	Score	Match	Length	DB	ID	Description
. 1	1986	100.0	1986		BD271616 X	BD271616 Novel cla
3	1986 1986	100.0	1,986 1986	6 6	BD271625 BD271626	BD271625 Novel cla BD271626 Novel cla

						AX037577 AX037595 AX037597 AX090382 SCYNR008W 4/96 SED PEUT SCN201952 DUP!	
	4	1986	100.0	1986	6	AX037577 QUP	AX037577 Sequence
	5	1986	100.0	1986	6	AX037595	AX037595 Sequence
	6	1986	100.0	1986	6	AX037597	AX037597 Sequence
	7.	1986	100.0	1986	6	AX090382 Juge SED Per	AX090382 Sequence
	8	1986	100'.0	2575	8	SCYNROO8W 4/10	Z71623 S.cerevisia
	9	1986	100.0	23901	8	SCN201952 DUP	X77395 S.cerevisia
С	10	26	1.3	135321	2	AC142021 火	AC142021 Rattus no
С	11	26	1.3	232722	2	AC112283 X	AC112283 Rattus no
С	12	23	1.2	203613	2	AC118575 X	AC118575 Lemur cat
С	13	22	1.1	173251	9	AL356280 X	AL356280 Human DNA
	14	22	1.1	283341	2	AC128766%	AC128766 Rattus no
С	15	21	1.1	1822	8	D88383 1997 Photograps	D88383 Hordeum vul
	16	21	1.1	1974	3	AF326965	AF326965 Trypanoso
	17	21	1.1	33744	9	AC027326	AC027326 Homo sapi
	18	21	1.1	78303	9	AC008910	AC008910 Homo sapi
	19	21		129575	9	AL160159	AL160159 Human DNA
	20	21	1.1	162349	9	AC006226	AC006226 Homo sapi
С	21	21		165518	3	AC091226	AC091226 Drosophil
	22	21		167345	5	BX004771	BX004771 Zebrafish
С	23	21		167.638	3	AC069405	AC069405 Drosophil
	24	21		188048	10	AL669974	AL669974 Mouse DNA
	25	21		193074	2	AC017576	AC017576 Drosophil
	26	21		199531	9	AC094104	AC094104 Homo sapi
	27.	21		200028	9	AC092680	AC092680 Homo sapi
	28	21		200229	10	AC099572	AC099572 Mus muscu
С	29	21		236957	2	AC118153	AC118153 Rattus no
	30	21		260198	2	BX469922	BX469922 Danio rer
С	31	21		282116	3	AE003537	AE003537 Drosophil
С	32	20	1.0	181	9	AY728770	AY728770 Macaca mu
С	33	20	1.0	450	6	CQ050097	CQ050097 Sequence
С	34	20	1.0	450	6	CQ065138	CQ065138 Sequence
С	35	20	1.0	450	6	CQ092094	CQ092094 Sequence
С	36	20	1.0	450	6	CQ130908	CQ130908 Sequence
С	37	20	1.0	450	6	CQ169518	CQ169518 Sequence
С	38	20	1.0	450	6	CQ198654	CQ198654 Sequence
С	39	20	1.0	450	6	CQ214119	CQ214119 Sequence
C	40	20	1.0	450	6	CQ252704	CQ252704 Sequence
Ċ	41	20	1.0	450	6	CQ289849	CQ289849 Sequence
С	42	20	1.0	450	6	CQ326815	CQ326815 Sequence
	43	20	1.0	459	8	AF072341	AF072341 Ajellomyc
	44	20	1.0	459	8	AF072342	AF072342 Ajellomyc
	45	20	1.0	459	8	AF072343	AF072343 Ajellomyc

OM nucleic - nucleic search, using sw model

Run on: May 2, 2005, 16:07:57; Search time 1102 Seconds

(without alignments)

10668.427 Million cell updates/sec

Title: US-09-937-779-1

Perfect score: 1986

Sequence: 1 atgggcacactgtttcgaag.....agatgcccttcccaatgtaa 1986

Scoring table: OLIGO NUC

Gapop 60.0 , Gapext 60.0

Searched: 4390206 seqs, 2959870667 residues

Word size : 0

Total number of hits satisfying chosen parameters: 8780412

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Listing first 45 summaries

Database : N_Geneseq_16Dec04:*

1: geneseqn1980s:*

2: geneseqn1990s:*

3: genesegn2000s:*

4: geneseqn2001as:*

5: genesegn2001bs:*

6: genesegn2002as:*

7: genesegn2002bs:*

8: geneseqn2003as:*

9: geneseqn2003bs:*

10: geneseqn2003cs:*

11: geneseqn2003ds:*

12: geneseqn2004as:*

13: geneseqn2004bs:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

Result No.	Score	Query Match	Length	DB	ID	Description
1	1986	100.0	1986	3	AAC64431 PUP	Aac64431 Saccharom
2	1986	100.0	1986		AAC64440	Aac64440 Saccharom
3	1986	100.0	1986	5	AAS01342	Aas01342 Yeast LCA
4	1986	100.0	1986	12	ADF47816 🗸	Adf47816 Yeast acv

	5	1694	85.3	1701	12	ADF47817 J D V AAC64441	Adf47817 Yeast mem
	6	1119	56.3	1986	3	AAC64441	Aac64441 Saccharom
	7	30	1.5	46	12	ADF47839 X	Adf47839 Yeast acy
	8	27	1.4	46	12	ADF47837 ×	Adf47837 Yeast acy
	9	25	1.3	41	12	ADF47838 X	Adf47838 Yeast acy
С	10	20	1.0	450	4	AAI11005 >	Aai11005 Probe #93
С	11	20	1.0	450	4	ABA52660 X	Aba52660 Human foe
С	12	20	1.0	450	4	AAI32267 X	Aai32267 Probe #95
С	13	20	1.0	450	4	ABA42236 ₹	Aba42236 Human bre
С	14	20	1.0	450	4	ABA22448 X	Aba22448 Probe #91
С	15	20	1.0	450	4	AAK26373	Aak26373 Human bon
С	16	20	1.0	450	4	AAK00918	Aak00918 Human bra
С	17	20	1.0	450	4	ABS25968	Abs25968 Human liv
С	18	20	1.0	450	5	AAI00926	Aai00926 Probe #91
С	19	20	1.0	450	6	ABS00963	Abs00963 Human gen
С	20	20	1.0	575	12		Ach68028 Human gen
С	21	20	1.0	792	5	AAF68302	Aaf68302 Human lun
С	22	20	1.0	792	6	ABK38213	Abk38213 cDNA enco
С	23	20	1.0	792	8	ACA10542	Acal0542 Human lun
С	24	20	1.0	792	8	ABX99493	Abx99493 Lung canc
С	25	20	1.0	792	10	ADH45739	Adh45739 Human lun
С	26	20	1.0	792	12	ADE72276	Ade72276 Human lun
С	27	20	1.0	792	13	ADJ19658	Adj19658 Human lun
	28	20	1.0	3420	10	ACF68206	Acf68206 Photorhab
С	29	20	1.0	110000	10	ACF67367 09	Continuation (10 o
С	30	20	1.0	110000	10	ACF65384 3	Continuation (4 of
	31	19	1.0	39	5	AAS01346	Aas01346 Yeast LRO
	32	19	1.0	540	4	AAK88546	Aak88546 Human dig
С	33	19	1.0	840	10	ADF02110	Adf02110 Bacterial
	34	19	1.0	951	5	AAH94468	Aah94468 Human foe
С	35	19	1.0	1056	13	ADS55204	Ads55204 Bacterial
С	36	19	1.0	1071	13	ADS49690	Ads49690 Bacterial
	37	19	1.0	1326	10	ACF71073	Acf71073 Photorhab
	38	19	1.0	1672	12	ADL13358	Adl13358 Human ste
	39	19	1.0	1819	4	AAK90612	Aak90612 Human dig
С	40	19	1.0	2000	8	ADA71449	Ada71449 Rice gene
С	41	19	1.0	2000	8	ADA72450	Ada72450 Rice gene
C	42	19	1.0	2000	12	ADJ41071	Adj41071 Plant cDN
C	43	19	1.0	2878	10	ADB68594	Adb68594 Mouse mic
	44	19	1.0	3176	13	ADR07994	Adr07994 Full leng
	45	19	1.0	3188	4	AAK90614	Aak90614 Human dig
							· 9

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OM nucleic - nucleic search, using sw model

Run on: May 2, 2005, 20:45:36; Search time 355 Seconds

(without alignments)

9153.931 Million cell updates/sec

Title: US-09-937-779-1

Perfect score: 1986

Sequence: 1 atgggcacactgtttcgaag.....agatgcccttcccaatgtaa 1986

Scoring table: OLIGO NUC

Gapop 60.0 , Gapext 60.0

Searched: 1202784 seqs, 818138359 residues

Word size: 0

Total number of hits satisfying chosen parameters: 2405568

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Listing first 45 summaries

1: /cgn2 6/ptodata/1/ina/5A COMB.seq:*

2: /cgn2 6/ptodata/1/ina/5B COMB.seq:*

3: /cgn2 6/ptodata/1/ina/6A COMB.seq:*

4: /cgn2_6/ptodata/1/ina/6B_COMB.seq:*

5: /cgn2 6/ptodata/1/ina/PCTUS COMB.seg:*

6: /cgn2 6/ptodata/1/ina/backfiles1.seq:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

			8	•					
Result			Query						
No.		Score	Match	Length	DB	ID	Description		
c	1	20	1.0	792	4	US-09-702-705-220 X	Sequence	220, A	
С	2	20	1.0	792	4	US-09-736-457-220 🗡	Sequence		
С	3	20	1.0	792	4	US-09-614-124B-220X	Sequence		pp
С	4	20	1.0	792	4	US-09-671-325-220 🏌	Sequence	220, A	pp
С	5	20	1.0	792	4	US-09-589-184-220	Sequence	220, A	pp
C	6	20	1.0	792	4	US-09-658-824-220 \	Sequence	220, A	pp
	7	19	1.0	601	4	US-09-949-016-72828	Sequence	72828,	A
	.8	19	1.0	601	4	US-09-949-016-72829	Sequence	72829,	Α
	9	19	1.0	601	4	US-09-949-016-72830	Sequence	72830,	Α
	10	19 '	1.0	601	4	US-09-949-016-72831	Sequence	72831,	Α
	11	19	1.0	601	4	US-09-949-016-72832 √	Sequence	72832,	Α

```
12
                                  US-09-949-016-72833×
            19
                  1.0
                          601
                                                                Sequence 72833, A
   13
            19
                  1.0
                          840
                               4
                                  US-09-543-681A-2395X
                                                                Sequence 2395, Ap
                                  US-09-248-796A-6352/99
   14
            19
                  1.0
                         1014
                               4
                                                                Sequence 635, App 6747137
   15
            19
                  1.0
                         1023
                                  US-09-248-796A-2948 1
                                                                Sequence 2948, Ap
   16
            19
                  1.0
                         1672
                                  US-09-976-594-1087
                                                                Sequence 1087, Ap
   17
            19
                  1.0
                         2878
                                  US-09-917-963-10
                                                                Sequence 10, Appl
С
   18
            19
                  1.0
                         4267
                                  US-09-661-753-47
                                                                Sequence 47, Appl
   19
С
            19
                  1.0 106450
                                  US-09-949-016-13873
                                                                Sequence 13873, A
   20
С
            19
                  1.0 147840
                                  US-09-949-016-15236
                                                                Sequence 15236, A
С
   21
            18
                  0.9
                           25
                                  US-09-396-196G-60894
                                                                Sequence 60894, A
С
   22
            18
                  0.9
                           36
                                  US-08-781-891-184
                                                                Sequence 184, App
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            18
                  0.9
                           36
                                  US-09-618-166-184
                                                                Sequence 184, App
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                          507
                                  US-09-248-796A-11763
                                                                Sequence 11763, A
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                                  US-09-949-016-115126
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                                                                Sequence 116421,
   27
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                               4
                                  US-09-949-016-116422
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                                  US-09-949-016-177105
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                                  US-09-949-016-203275
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                         2090
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                         2106
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С
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                                  US-09-815-923-13
                                                                Sequence 13, Appl
С
   36
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                         9060
                               3
                                  US-08-378-313-20
                                                                Sequence 20, Appl
   37
            18
                  0.9
                       19703
                                  US-09-949-016-14926
                                                                Sequence 14926, A
   38
           18
                  0.9
                       25512
                                  US-09-949-016-15886
                                                                Sequence 15886, A
   39
           18
                  0.9
                       25512
                               4
                                  US-09-949-016-15887
                                                                Sequence 15887, A
   40
                                  US-09-949-016-17478
           18
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                        49472
                               4
                                                                Sequence 17478, A
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           18
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                       50878
                               4
                                  US-09-949-016-11787
                                                                Sequence 11787, A
   42
                  0.9
           18
                       92344
                                  US-09-949-016-16802
                                                                Sequence 16802, A
   43
            18
                  0.9
                       93364
                                  US-09-949-016-14890
                                                                Sequence 14890, A
   44
            18
                  0.9 101349
                                  US-09-949-016-17433
                                                                Sequence 17433, A
   45
           18
                  0.9 173787
                                  US-09-949-016-12542
                                                                Sequence 12542, A
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OM nucleic - nucleic search, using sw model

Run on: May 2, 2005, 20:51:36; Search time 1200 Seconds

(without alignments)

10075.121 Million cell updates/sec

Title: US-09-937-779-1

Perfect score:

Sequence: 1 atgggcacactgtttcgaag.....agatgcccttcccaatgtaa 1986

Scoring table: OLIGO NUC

Gapop 60.0 , Gapext 60.0

5642217 seqs, 3043843248 residues Searched:

Word size :

Total number of hits satisfying chosen parameters: 11284434

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Listing first 45 summaries

Database : Published Applications NA:*

/cgn2_6/ptodata/1/pubpna/US07 PUBCOMB.seq:* 1:

2: /cgn2_6/ptodata/1/pubpna/PCT NEW PUB.seq:*

3: /cgn2_6/ptodata/1/pubpna/US06 NEW PUB.seq:*

/cgn2_6/ptodata/1/pubpna/US06_PUBCOMB.seq:*

/cgn2 6/ptodata/1/pubpna/US07 NEW PUB.seg:*

/cgn2_6/ptodata/1/pubpna/PCTUS_PUBCOMB.seq:*

/cgn2_6/ptodata/1/pubpna/US08_NEW_PUB.seq:* 7:

/cgn2 6/ptodata/1/pubpna/US08 PUBCOMB.seq:*

/cgn2 6/ptodata/1/pubpna/US09A PUBCOMB.seg:*

10: /cgn2 6/ptodata/1/pubpna/US09B PUBCOMB.seg:*

11:

/cgn2_6/ptodata/1/pubpna/US09C_PUBCOMB.seq:* 12:

/cgn2_6/ptodata/1/pubpna/US09_NEW_PUB.seq:* 13:

/cgn2_6/ptodata/1/pubpna/US10A PUBCOMB.seq:* 14: /cgn2_6/ptodata/1/pubpna/US10B PUBCOMB.seq:*

15: /cgn2_6/ptodata/1/pubpna/US10C_PUBCOMB.seq:*

/cgn2 6/ptodata/1/pubpna/US10D PUBCOMB.seg:*

17:

/cgn2 6/ptodata/1/pubpna/US10E PUBCOMB.seq:*

/cgn2_6/ptodata/1/pubpna/US10F PUBCOMB.seq:*

/cgn2_6/ptodata/1/pubpna/US10_NEW_PUB.seq:* 19:

20: /cgn2 6/ptodata/1/pubpna/US11_NEW_PUB.seq:*

/cgn2_6/ptodata/1/pubpna/US60_NEW_PUB.seq:* 21:

/cgn2 6/ptodata/1/pubpna/US60 PUBCOMB.seg:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

Res	111t		* Query				
	No.	Score	-	Length	פח	ID	Doggrintion
				nengen			Description
	1	21	1.1	183334	19	US-10-741-600-17646 X	Sequence 17646, A
С	2	20	1.0	367	18	US-10-425-115-147325.	Sequence 147325,
С	3	20	1.0	450	9	US-09-864-761-914	Sequence 914, App
c	4	20	1.0	575	16	US-10-029-386-1223	Sequence 1223, Ap
c	5	. 20	1.0	792	9	US-09-736-457-220	Sequence 220, App
c	6	20	1.0	792	9	US-09-902-941-220	Sequence 220, App
c	7	20	1.0	792	9	US-09-849-626-220	Sequence 220, App
c	8	20	1.0	792	10	US-09-476-300-220	Sequence 220, App
c	9	20	1.0	792	14	US-10-017-754-220	
c	10	20	1.0	792	16	US-10-113-872-220	Sequence 220, App Sequence 220, App
c	11	20	1.0	792	17	US-10-283-017-220	Sequence 220, App
·	12	20	1.0	834	18	US-10-437-963-91894	Sequence 91894, A
·c	13	20	1.0		17	ÚS-10-424-599-109347	
c	14	20	1.0	3089	18	US-10-425-115-109147	Sequence 109347,
C	15	20	1.0	4833	18	US-10-425-115-109147-	Sequence 109147,
•	16	20	1.0	4950	18	US-10-425-115-177412	Sequence 177412,
С	17	20	1.0	5433	18	US-10-425-115-109132 US-10-425-115-177411	Sequence 109152,
c	18	19	1.0	25	19	US-10-719-900-547596	Sequence 177411,
c	19	19	1.0	201	18	US-10-719-993-17098	Sequence 547596,
C	20	19	1.0	201	18	US-10-719-993-17098 US-10-719-993-43317	Sequence 17098, A
	21	19	1.0	415	18		Sequence 43317, A
С	22	19	1.0	740	18	US-10-437-963-15580 US-10-437-963-67956	Sequence 15580, A
c	23	19	1.0	1056	17		Sequence 67956, A
С	24	19	1.0	1036	17	US-10-369-493-30878	Sequence 30878, A
C	25	19	1.0	1930	17	US-10-369-493-28120	Sequence 28120, A
•	26	19	1.0	2000	17	US-10-425-114-26751	Sequence 26751, A
c c	27	19	1.0	2878		US-10-260-238-2071	Sequence 2071, Ap
c	28	19			10	US-09-917-963-10	Sequence 10, Appl
C	29	19	1.0	2878	19	US-10-764-420-2702	Sequence 2702, Ap
_	30		1.0	3007	18	US-10-425-115-173430	Sequence 173430,
С	31	19 19	1.0	. 3192 3822	18	US-10-425-115-87876	Sequence 87876, A
	32	19	1.0	3864	18	US-10-664-705-148	Sequence 148, App
_	33	19	1.0		17	US-10-369-493-25515	Sequence 25515, A
C C	34	19	1.0	4267 4267	10	US-09-948-002-47	Sequence 47, Appl
С	35	19		4267	17	US-10-189-267-11	Sequence 11, Appl
C	36	19	1.0 1.0	5218	17 17	US-10-633-163-47	Sequence 47, Appl
	37	19	1.0	10006		US-10-310-154-250	Sequence 250, App
	38	19	1.0	13051	15	US-10-311-455-9	Sequence 9, Appli
_	39	19	1.0	50287	18	US-10-719-993-6932	Sequence 6932, Ap
C C	40	19		107829	13 13	US-10-087-192-1501	Sequence 1501, Ap
C	41	19		168407	18	US-10-087-192-361	Sequence 361, App
_	42	19		366803	18	US-10-322-281-305	Sequence 305, App
C C	43	18	0.9	25		US-10-719-993-6805	Sequence 6805, Ap
c	44	18	0.9	36	19 17	US-10-809-189-60894 US-10-374-077-184	Sequence 60894, A
C	45	18	0.9	260	11	US-09-922-293-1957	Sequence 184, App
	30	10	0.5	200	ΤT	03203-322-233-1337	Sequence 1957, Ap

OM nucleic - nucleic search, using sw model

Run on: May 2, 2005, 20:25:31; Search time 6685 Seconds

(without alignments)

11308.248 Million cell updates/sec

Title: US-09-937-779-1

Perfect score: 1986

Sequence: 1 atgggcacactgtttcgaag.....agatgcccttcccaatgtaa 1986

Scoring table: OLIGO NUC

Gapop 60.0 , Gapext 60.0

Searched: 34239544 seqs, 19032134700 residues

Word size: 0

Total number of hits satisfying chosen parameters: 68479088

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Listing first 45 summaries

Database : EST:*

1: gb_est1:*

2: gb_est2:*

3: gb_htc:*

4: gb_est3:*

5: gb_est4:*

6: gb_est5:*

7: gb_est6:*

8: gb_gssl:*

9: gb_gss2:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

Resi	ılt		% Query					
	No.	Score		Length	DB	ID	Descript	Lon
c	1	23	1.2	400	8	AZ924376 DUP	AZ924376	4906.ic28
	2	22	1.1	553	8	AZ525082 X	AZ525082	240PbA04
С	3	22	1.1	564	6	CD847546 X	CD847546	DHOAB60ZH
С	4	. 22	1.1	620	7	CF077965 \	CF077965	QHK17N06.
С	5	22	1.1	752	5	BU026006		OHG12L10.
	6	21	1.1	181	8	ВН738251	BH738251	BOHXI49TR
	7	21	1.1	183	7	CF918634	CF918634	Bflor498.
	8	21	1.1	184	7	CF918690 √	CF918690	Bflor498.

*										
	9	21	1.1	256	7 .	CN191467		CN191467	HCBCSO6 O	
С	10	21	1.1	288	4	BJ454259		BJ454259		
c	11	21	1.1	288	9	CL206068		CL206068		
	12	21	1.1	338	8	BH440971		BH440971		
С	13	21	1.1	386	4	BM443004		BM443004		
	14	21	1.1	393	8	V.		BH690705		
	15	21	1.1	403	8	B03865 1990	prm	B03865 cSi		
	16	21	1.1		4	BJ461791	prmers	BJ461791		
	17	21	1.1	457	4	BI385211		BI385211		
С	18	21	1.1	461	9	CL245339		CL245339		
	19	21	1.1	464	8	BH439796		BH439796		
	20	21	1.1	485	4	BI385212		BI385212		
	21	21	1.1	537	8	BH461140		BH461140 I		
	22	21	1.1	554	8	BH590618		BH590618 1		
	23	21	1.1	573	8	AQ778682		AQ778682		
С	24	21	1.1	596	8	BH669448		BH669448 1		
С	25	21	1.1	597	6	CD055309		CD055309 I	HO08N23S	
	26	21	1.1	598	9	CC948120		CC948120 I	BOIBR05TF	
	27	21	1.1	601	8	BH650596		BH650596 1	BOHYC18TR	
C	28	21	1.1	605	8	BH720093		BH720093 1	BOMLN41TR	
	29	21	1.1	607	8	BH455709		BH455709 1	BOHSH59TF	
С	30	21	1.1	622	5	BQ793370		BQ793370 1	EST 2308	
	31	21	1.1	633	7	CK403317		CK403317	AUF_IfHdk	
	32	21	1.1	639	8	BH424160		BH424160 I	BOGTH82TR	
С	33	21	1.1	639	9	CG737180	-	CG737180	ZMMBBb032	
	34	21	1.1	642	8	BH682774		BH682774 I		
С	35	21	1.1	649	5	BQ793616		BQ793616 I		
С	36	21	1.1	664	9	CC968690		CC968690 I		
C	37	21	1.1	667	8	BH250391		BH250391		
	38	21	1.1	668	8	BH688265	• •	BH688265 1		
	39	21	1.1	680	8	BH972432		BH972432		
	40	21	1.1	680	8	BZ072897		BZ072897 .		,
С	41	21	1.1	688	8	AQ953709		AQ953709		
	42	21	1.1	694	8	BZ047486		BZ047486		
	43	21	1.1	695	8	BH541538		BH541538 I		
	44	21	1.1	696	8	BH734523		BH734523 1		
	45	21	1.1	696	8	вн930691		вн930691	odj01e06.	

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OM protein - nucleic search, using frame plus p2n model Run on: May 2, 2005, 11:27:24; Search time 7744 Seconds (without alignments) 4135.963 Million cell updates/sec Title: US-09-937-779-2 Perfect score: Sequence: 1 MGTLFRRNVQNQKSDSDENN......VEPRQLSNLSQWVSQMPFPM 661 Scoring table: BLOSUM62 Xgapop 10.0 , Xgapext 0.5 Ygapop 10.0 , Ygapext 0.5 Fgapop 6.0 , Fgapext 7.0 Delop 6.0 , Delext 7.0 Searched: 4708233 segs, 24227607955 residues Total number of hits satisfying chosen parameters: 9416466 Minimum DB seq length: 0 Maximum DB seq length: 2000000000 Post-processing: Minimum Match 0% Maximum Match 100% Listing first 45 summaries Command line parameters: -MODEL=frame+ p2n.model -DEV=xlh Q=/cgn2_1/USPTO_spool/US09937779/runat_02052005_122715_18261/app query.fasta 1.8 -DB=GenEmbl -QFMT=fastap -SUFFIX=rge -MINMATCH=0.1 -LOOPCL=0 -LOOPEXT=0 -UNITS=bits -START=1 -END=-1 -MATRIX=blosum62 -TRANS=human40.cdi -LIST=45 -DOCALIGN=200 -THR SCORE=pct -THR MAX=100 -THR MIN=0 -ALIGN=15 -MODE=LOCAL -OUTFMT=pto -NORM=ext -HEAPSIZE=500 -MINLEN=0 -MAXLEN=2000000000 -USER=US09937779_@CGN_1_1_4545 @runat 02052005 122715 18261 -NCPU=6 -ICPU=3 -NO MMAP -LARGEQUERY -NEG SCORES=0 -WAIT -DSPBLOCK=100 -LONGLOG -DEV TIMEOUT=120 -WARN TIMEOUT=30 -THREADS=1 -XGAPOP=10 -XGAPEXT=0.5 -FGAPOP=6 -FGAPEXT=7 -YGAPOP=10 -YGAPEXT=0.5 -DELOP=6 -DELEXT=7 Database : GenEmbl: * 1: gb ba:* gb htg:* ' 3: gb in: * gb om: * 4: 5: gb_ov:* 6: gb pat:* 7: gb ph:* gb_pl:* 8:

9:

gb pr:* 10: gb ro:* 11: gb_sts:*
12: gb_sy:*
13: gb_un:*
14: gb_vi:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

			0			SUPPLAKTES	
ъ	1 .		8			•	
Res		_	Query				
	No.	Score	Match	Length	DB	ID	Description
	1	3546	100.0	1986	6	BD271616 DUD (BD271616 Novel cla
	2	3546	100.0	1986	6	BD271625	BD271625 Novel cla
	3	3546	100.0	1986	6	BD271626	BD271626 Novel cla
	4	3546	100.0	1986	6	AX037577	AX037577 Sequence
	5	3546	100.0	1986	6	AX037595	AX037595 Sequence
	6	3546	100.0	1986	6	AX037597	AX037597 Sequence
	7	3546	100.0	1986	6	AX090382	AX090382 Sequence
•	8	3546	100.0	2575	8	SCYNR008W	Z71623 S.cerevisia
	9	3546	100.0	23901	8	SCN201952	X77395 S.cerevisia
c ·	10	2469		110000	8	CR380955 05	Continuation (6 of
С	11	2244.5		110000	8	AE016819 00	AE016819 Eremothec
c	12	2244.5		110000	8	AE016819 01	Continuation (2 of
c	13	2220.5		110000	8	CR382126 06	Continuation (7 of
_	14	1840		110000	8	CR382136 07	Continuation (8 of
	15	1840		110000	8	CR382136 08	Continuation (9 of
	16	1769	49.9	2106	6	AR546016	AR546016 Sequence
_	17	1630		110000	8	CR382131 19	Continuation (20 o
. С	18	1245.5		103568	8	CNS07EGJ	·
•	19	1245.5					AL590462 DNA centr
С				322194	8	CNS09S4S	BX088700 DNA centr
	20	1150.5	32.4	2312	6	BD271617	BD271617 Novel cla
	21	1150.5	32.4	2312	6	BD271627	BD271627 Novel cla
	22	1150.5	32.4	2312	6	AX037579	AX037579 Sequence
	23	1150.5	32.4	2312	6	AX037599	AX037599 Sequence
	24	1150.5	32.4	42391	8	SPBC776	AL035263 S.pombe c
	25	852	24.0	4093	6	AX794722	AX794722 Sequence
	26	845.5	23.8	2828	8	AY210981	AY210981 Medicago
	27	839.5	23.7	2433	6	AX794708	AX794708 Sequence
	28	839.5	23.7	2495	8	AK100079	AK100079 Oryza sat
	29	839.5	23.7	2565	6	AX794706	AX794706 Sequence
	30	836.5	23.6	2479	6	AX794704	AX794704 Sequence
	31	835	23.5	2700	6	AX794710	AX794710 Sequence
	32	832.5	23.5	2441	8	BT013705	BT013705 Lycopersi
	33	832	23.5	2013	6	AX090380	AX090380 Sequence
	34	832	23.5	2016	6	AX412864	AX412864 Sequence
	35	832	23.5	2402	8	AY052715	AY052715 Arabidops
	36	. 832	23.5	2425	6	AX925713	AX925713 Sequence
	37	828	23.4	2398	6	AX794712	AX794712 Sequence
	38	823	23.2	2427	6	BD271619	BD271619 Novel cla
	39	823	23.2	2427	6	AX037581	AX037581 Sequence
	40	823	23.2	2431	8	AY160110	AY160110 Arabidops
	41	819	23.1	2030	6	AX794714	AX794714 Sequence
С	42	758.5	21.4	39475	2	AC149358	AC149358 Phakopsor
Ŭ	43	724	20.4	3203	8	AK099811	AK099811 Oryza sat
		,24	20.4	5205	U	AMOJJUTI	MRUJJOII OLYZA SAL

OM protein - nucleic search, using frame plus p2n model May 2, 2005, 11:27:24; Search time 1036 Seconds Run on: (without alignments) 3776.978 Million cell updates/sec Title: US-09-937-779-2 Perfect score: 3546 Sequence: 1 MGTLFRRNVQNQKSDSDENN......VEPRQLSNLSQWVSQMPFPM 661 Scoring table: BLOSUM62 Xgapop 10.0 , Xgapext 0.5 Ygapop 10.0 , Ygapext 0.5 Fgapop 6.0 , Fgapext 7.0 Delop 6.0 , Delext 7.0 4390206 segs, 2959870667 residues Searched: Total number of hits satisfying chosen parameters: 8780412 Minimum DB seg length: 0 Maximum DB seq length: 2000000000 Post-processing: Minimum Match 0% Maximum Match 100% Listing first 45 summaries Command line parameters: -MODEL=frame+ p2n.model -DEV=x1h Q=/cgn2_1/USPTO_spool/US09937779/runat_02052005_122714_18249/app_query.fasta_1.8 -DB=N Geneseq 16Dec04 -QFMT=fastap -SUFFIX=rng -MINMATCH=0.1 -LOOPCL=0 -LOOPEXT=0 -UNITS=bits -START=1 -END=-1 -MATRIX=blosum62 -TRANS=human40.cdi -LIST=45 -DOCALIGN=200 -THR SCORE=pct -THR MAX=100 -THR MIN=0 -ALIGN=15 -MODE=LOCAL -OUTFMT=pto -NORM=ext -HEAPSIZE=500 -MINLEN=0 -MAXLEN=2000000000 -USER=US09937779_@CGN_1_1_470_@runat_02052005_122714_18249 -NCPU=6 -ICPU=3 -NO MMAP -LARGEQUERY -NEG SCORES=0 -WAIT -DSPBLOCK=100 -LONGLOG -DEV_TIMEOUT=120 -WARN_TIMEOUT=30 -THREADS=1 -XGAPOP=10 -XGAPEXT=0.5 -FGAPOP=6 -FGAPEXT=7 -YGAPOP=10 -YGAPEXT=0.5 -DELOP=6 -DELEXT=7 Database : N Geneseq 16Dec04:* 1: genesegn1980s:* geneseqn1990s:* 2: 3: geneseqn2000s:* 4: geneseqn2001as:* 5: geneseqn2001bs:* 6: geneseqn2002as:* 7: geneseqn2002bs:* geneseqn2003as:*

9:

geneseqn2003bs:* 10: geneseqn2003cs:* 11: geneseqn2003ds:*
12: geneseqn2004as:*
13: geneseqn2004bs:*

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Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

Res	sult		Query				
	No.	Score		Length	DB	ID	Description
						•	
	1	3546	100.0	. 1986	3	AAC64431 Dupl	Aac64431 Saccharom
	2	3546	100.0	1986	3	AAC64440	Aac64440 Saccharom
	3	3546	100.0	1986	5	AAS01342	Aas01342 Yeast LCA
	4	3546	100.0	1986	12	ADF47816 ·	Adf47816 Yeast acy
	5	3481	98.2	1986	3	AAC64441	Aac64441 Saccharom
	6	3029	85.4	1701	. 12	ADF47817 /	Adf47817 Yeast mem
	7	1269	35.8	1872	12	ADF47831 🌱	Adf47831 Schizosac
С	8	1182	33.3	7299	13	ADR84383	Adr84383 Aspergill
	9	1165.5	32.9	2223	12	ADF47830	Adf47830 Aspergill
	10	1155	32.6	2047	12	ADF47829	Adf47829 Aspergill
	11	1150.5	32.4	2312	3	AAC64432 /	Aac64432 Schizosac
	12	1132.5	31.9	2312	3	AAC64442 /	Aac64442 Schizosac
	13	852	24.0	4093	9	AAL62913 💢	Aal62913 Soybean c
	14	839.5	23.7	2433	9	AAL62907 2001	Aa162907 Rice cDNA
	15	839.5	23.7	2565	9	AAL62906 X	Aa162906 Corn cDNA
	16	838.5	23.6	2004	12	ADF47835	Adf47835 Crepis pa
	1,7	836.5	23.6	2479	9	AAL62905	Aa162905 Guayule c
	18	835	23.5	2700	9	AAL62908	Aa162908 Soybean c
	19	832	23.5	2013	5	AAS01341	Aas01341 Arabidops
	20	832	23.5	2016	6	ADG88186	Adg88186 A. thalia
	21	832	23.5	2016	12	ADF47819	Adf47819 Arabidops
	22	832	23.5	2425	10	ADG25155	Adg25155 P. patens
	23	828	23.4	2398	9	AAL62909	Aa162909 Sunflower
	24	823	23.2	2427	3	AAC64434 ·	Aac64434 Arabidops
	25	822.5	23.2	1803	12	ADF47821	Adf47821 Arabidops
	26	819	23.1	2030	9	AAL62910	Aa162910 Wheat cDN
	27	807.5	22.8	1998	8	ABZ76360	Abz76360 A. thalia
	28	807.5	22.8	1998	. 12	ADF47827	Adf47827 Arabidops
	29	806.5	22.7	2016	12	ADF47833	Adf47833 Crepis pa
	30	539	15.2	3107	5	AAS01085	Aas01085 Arabidops
	31	532	15.0	3685	3	AAC64433	Aac64433 Arabidops
	32	526	14.8	3685	3	AAC64443	Aac64443 Arabidops
	33	457.5	12.9	553	8	ABZ54921	Abz54921 Aspergill
	34	410	11.6	578	13	ADR61378	Adr61378 Cotton cD
	35	403	11.4	1510	9	AAL62904	Aal62904 Guar cDNA
С	36	320	9.0	1680	5	AAS01086	Aas01086 Arabidops
	37	309	8.7	616	3	AAC64436	Aac64436 Neurospor
	38	301.5	8.5	1902	12	ADF47823	Adf47823 Arabidops
	39	299.5	8.4	1641	5	AAS01082	Aas01082 Arabidops
	40	254.5	7.2	1323	6	ABK87900	Abk87900 cDNA enco
	41	254.5	7.2	1358	1	AAN70191	Aan70191 DNA clone
	42	254.5	7.2	1744	2	AAZ32180	Aaz32180 Human lec
	43	254.5	7.2	1744	5	AAD02648	Aad02648 Human lec
	44	254.5	7.2	1744	6	ABK35510	Abk35510 Human end

OM protein - nucleic search, using frame_plus_p2n model

Run on: May 2, 2005, 11:27:24; Search time 306 Seconds

(without alignments)

3534.572 Million cell updates/sec

Title: US-09-937-779-2

Perfect score: 3546

Sequence: 1 MGTLFRRNVQNQKSDSDENN.....VEPRQLSNLSQWVSQMPFPM 661

Scoring table: BLOSUM62

Xgapop 10.0 , Xgapext 0.5 Ygapop 10.0 , Ygapext 0.5 Fgapop 6.0 , Fgapext 7.0 Delop 6.0 , Delext 7.0

Searched: 1202784 segs, 818138359 residues

Total number of hits satisfying chosen parameters: 2405568

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Command line parameters:

-MODEL=frame+ p2n.model -DEV=x1h

Q=/cgn2_1/USPTO_spoo1/US09937779/runat_02052005_122715_18282/app_query.fasta_1.8

-DB=Issued Patents NA -QFMT=fastap -SUFFIX=rni -MINMATCH=0.1 -LOOPCL=0

-LOOPEXT=0 -UNITS=bits -START=1 -END=-1 -MATRIX=blosum62 -TRANS=human40.cdi

-LIST=45 -DOCALIGN=200 -THR SCORE=pct -THR MAX=100 -THR MIN=0 -ALIGN=15

-MODE=LOCAL -OUTFMT=pto -NORM=ext -HEAPSIZE=500 -MINLEN=0 -MAXLEN=2000000000

-USER=US09937779 @CGN 1 1 69 @runat_02052005_122715_18282 -NCPU=6 -ICPU=3

-NO MMAP -LARGEQUERY -NEG SCORES=0 -WAIT -DSPBLOCK=100 -LONGLOG

-DEV_TIMEOUT=120 -WARN_TIMEOUT=30 -THREADS=1 -XGAPOP=10 -XGAPEXT=0.5 -FGAPOP=6

-FGAPEXT=7 -YGAPOP=10 -YGAPEXT=0.5 -DELOP=6 -DELEXT=7

Database : Issued Patents NA:*

1: /cgn2 6/ptodata/1/ina/5A COMB.seq:*

2: /cgn2 6/ptodata/1/ina/5B COMB.seq:*

3: /cgn2_6/ptodata/1/ina/6A_COMB.seq:*

4: /cgn2 6/ptodata/1/ina/6B COMB.seq:*

5: /cgn2 6/ptodata/1/ina/PCTUS COMB.seq:*

6: /cgn2 6/ptodata/1/ina/backfiles1.seg:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

Resu	11 t	•	% Query				
	No.	Score	-	Length	DB	ID	Description
		1760					
	1	1769	49.9			US-09-248-796A-1147 PVD	Sequence 1147, Ap
	2	254.5	7.2		4	US-09-949-016-3703 X	Sequence 3703, Ap 66927,728 Sequence 33, Appl Lander, 728 Sequence 30, Appl 6
	3	254.5	7.2		4	US-09-054-272-33 PD4198	Sequence 33, Appl Lander 1022
	4	254.5	7.2		4	US-09-919-497-30 X	Sequence 30, Appl 6
	5	246.5	7.0	1236	4	US-09-402-532-17 /990	Sequence 17, Appl6498019 1026
	6	245.5	6.9	1332	4	US-09-402-532-18 //	Sequence 18, Appl 11
	7	244.5	6.9	1137	4	US-09-402-532-12 μ	Sequence 12, Appl //
	8	244.5	6.9	1146	4	US-09-402-532-15	Sequence 15, Appl
	9	243.5	6.9	1233	4	US-09-402-532-13	Sequence 13, Appl
	10	243.5	6.9		4	US-09-402-532-16	Sequence 16, Appl
	11	242.5	6.8	1236	4	US-09-402-532-19	Sequence 19, Appl
	12	237.5	6.7		4	US-09-402-532-14 V	Sequence 14, Appl
	13	233.5	6.6	2687	4	US-09-489-847-57 [998	Sequence 57, Appl 6476195/08
С	14	127	3.6	2640	4	US-09-252-991A-11547/990	Sequence 11547, A
	15	126	3.6	2934	4	US-09-252-991A-11690/948	Sequence 11690, A 3 7 725
	16	123	3.5	8230	4	US-09-949-016-15445	Sequence 15445, A
	17	122	3.4	552	4	US-09-248-796A-10377	Sequence 10377, A
С	18	122	3.4	2307	4	US-09-252-991A-6437	Sequence 6437, Ap
	19	120	3.4	7430	4	US-08-956-171E-260	Sequence 260, App
	20	120	3.4	7430	4	US-08-781-986A-260	Sequence 260, App
	21	115.5	3.3	2289	4	US-09-107-532A-621	Sequence 621, App
	22	114.5	3.2	2032	4	US-09-949-016-2678	Sequence 2678, Ap
	23	114.5	3.2	2032	4	US-09-949-016-2700	Sequence 2700, Ap
	24	114.5	3.2	6032	4	US-09-949-016-14420	Sequence 14420, A
	25	114.5	3.2	6033	4	US-09-949-016-14442	Sequence 14442, A
	26	113.5	3.2	1098	4	US-09-248-796A-6883	Sequence 6883, Ap
	27	111	3.1	2226	4	US-09-252-991A-14877	Sequence 14877, A
С	28	111	3.1	2268	4	US-09-252-991A-14487	Sequence 14487, A
	29	110.5	3.1	2038	4	US-09-949-016-1379	Sequence 1379, Ap
	30	110	3.1	3114	4	US-09-543-681A-3505	Sequence 3505, Ap
С	31	109.5		1664976		US-08-916-421B-1	Sequence 1, Appli
С	32	109.5		1664976	4		Sequence 1, Appli
	33	107	3.0	4404	4	US-09-134-000C-3335	Sequence 3335, Ap .
	34	105	3.0	1380	4	US-09-252-991A-6030	Sequence 6030, Ap
	35	105	3.0	1530	4	US-09-252-991A-6116	Sequence 6116, Ap
	36	105	3.0	1611	4	US-09-600-099-2	Sequence 2, Appli
	37	105	3.0		4	US-09-600-099-1	Sequence 1, Appli
C	38	104.5	2.9		3	US-08-961-527-89	Sequence 89, Appl
	39	104.5		129908	4	US-09-585-858-1	Sequence 1, Appli
	40	104.5		129908	4	US-10-270-878-1	Sequence 1, Appli
С	41	103	2.9	4403765	3	US-09-103-840A-2	Sequence 2, Appli
С	42	103		4411529	3	US-09-103-840A-1	Sequence 1, Appli
	43	102.5	2.9	2169	3	US-08-806-326-5	Sequence 5, Appli
С	44	102	2.9		4	US-09-902-540-1214	Sequence 1214, Ap
	45	101	2.8	1077	3	US-09-155-920-1	Sequence 1, Appli

OM protein - nucleic search, using frame plus p2n model Run on: May 2, 2005, 11:27:24; Search time 997 Seconds (without alignments) 4036.069 Million cell updates/sec Title: US-09-937-779-2 Perfect score: 3546 1 MGTLFRRNVQNQKSDSDENN......VEPRQLSNLSQWVSQMPFPM 661 Sequence: Scoring table: BLOSUM62 Xgapop 10.0 , Xgapext 0.5 Ygapop 10.0 , Ygapext 0.5 Fgapop 6.0 , Fgapext 7.0 Delop 6.0 , Delext 7.0 5642217 seqs, 3043843248 residues Searched: Total number of hits satisfying chosen parameters: 11284434 Minimum DB seq length: 0 Maximum DB seq length: 2000000000 Post-processing: Minimum Match 0% Maximum Match 100% Listing first 45 summaries Command line parameters: -MODEL=frame+ p2n.model -DEV=x1h Q=/cgn2 1/USPTO spool/US09937779/runat 02052005 122715 18296/app query.fasta 1.8 -DB=Published Applications NA -QFMT=fastap -SUFFIX=rnpb -MINMATCH=0.1 -LOOPCL=0 -LOOPEXT=0 -UNITS=bits -START=1 -END=-1 -MATRIX=blosum62 -TRANS=human40.cdi -LIST=45 -DOCALIGN=200 -THR SCORE=pct -THR MAX=100 -THR_MIN=0 -ALIGN=15 -MODE=LOCAL -OUTFMT=pto -NORM=ext -HEAPSIZE=500 -MINLEN=0 -MAXLEN=2000000000 -USER=US09937779 @CGN 1 1 480 @runat 02052005 122715 18296 -NCPU=6 -ICPU=3 -NO_MMAP -LARGEQUERY -NEG_SCORES=0 -WAIT -DSPBLOCK=100 -LONGLOG -DEV_TIMEOUT=120 -WARN_TIMEOUT=30 -THREADS=1 -XGAPOP=10 -XGAPEXT=0.5 -FGAPOP=6 -FGAPEXT=7 -YGAPOP=10 -YGAPEXT=0.5 -DELOP=6 -DELEXT=7 Database : Published Applications NA:* 1: /cgn2_6/ptodata/1/pubpna/US07_PUBCOMB.seq:* 2: /cgn2_6/ptodata/1/pubpna/PCT NEW PUB.seq:* 3: /cgn2_6/ptodata/1/pubpna/US06 NEW PUB.seq:* /cgn2_6/ptodata/1/pubpna/US06_PUBCOMB.seq:* /cgn2 6/ptodata/1/pubpna/US07 NEW PUB.seq:* /cgn2 6/ptodata/1/pubpna/PCTUS PUBCOMB.seq:* /cgn2 6/ptodata/1/pubpna/US08 NEW PUB.seq:* /cgn2_6/ptodata/1/pubpna/US08_PUBCOMB.seq:*

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13:
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14:
     /cgn2_6/ptodata/1/pubpna/US10B PUBCOMB.seq: *
15:
     /cgn2 6/ptodata/1/pubpna/US10C PUBCOMB.seq:*
16:
     /cgn2 6/ptodata/1/pubpna/US10D PUBCOMB.seg:*
17:
     /cgn2_6/ptodata/1/pubpna/US10E_PUBCOMB.seq:*
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19:
     /cgn2_6/ptodata/1/pubpna/US10_NEW_PUB.seq:*
20:
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21:
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     /cgn2_6/ptodata/1/pubpna/US60 PUBCOMB.seq:*
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Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

욯

Result No.		Caoro	Query	Tonath	DB	TD	
		Score	Match	Length		ID	Description .
	1	1630	46.0	2326	19	US-10-883-760-45 Pup/	Sequence 45, Appl
	2	852	24.0	4093	17	US-10-321-802-35 X	Sequence 35, Appl
	3	839.5	23.7	2433	17	US-10-321-802-21 \	Sequence 21, Appl
	4	839.5	23.7	2565	17	US-10-321-802-19	Sequence 19, Appl
	5	836.5	23.6	2479	17	US-10-321-802-17	Sequence 17, Appl
С	6	835	23.5	2395	18	US-10-437-963-38278	Sequence 38278, A
	7	835	23.5	2700	17	US-10-321-802-23	Sequence 23, Appl
	8	828	23.4	2398	17	US-10-321-802-25	Sequence 25, Appl
	9	824.5	23.3	2700	18	US-10-425-115-55049	Sequence 55049, A
	10	819	23.1	2030	17	US-10-321-802-27	Sequence 27, Appl
	11	814	23.0	2838	17	US-10-424-599-12571B	Sequence 125713,
	12	807.5	22.8	1998	16	US-10-217-939-5 ✓	Sequence 5, Appli
C	13	576.5	16.3	1388	18	US-10-425-115-55048	Sequence 55048, A
	14	410	11.6	578	18	US-10-767-795-2159	Sequence 2159, Ap
	15	403	11.4	1510	17	US-10-321-802-15	Sequence 15, Appl
	16	357.5	10.1	735	18	US-10-437-963-38267	Sequence 38267, A
	17	298	8.4	1141	17	US-10-424-599-124418	Sequence 124418,
	18	293	8.3	887	18	US-10-767-701-9911	Sequence 9911, Ap
	19	254.5	7.2	1744	9	US-09-919-497-30	Sequence 30, Appl
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	21	246.5	7.0	1236	16	US-10-323-051-17	Sequence 17, Appl
	22	246.5	7.0	1400	18	US-10-755-889-577	Sequence 577, App
	23	246.5	7.0	2680	9	US-09-978-295A-156	Sequence 156, App
	24	246.5	7.0	2680	9	US-09-978-697-156	Sequence 156, App
	25	246.5	7.0	2680	9	US-09-978-192A-156	Sequence 156, App
	26	246.5	7.0	2680	9	US-09-999-832A-156	Sequence 156, App
	27	246.5	7.0	2680	10	US-09-978-189-156	Sequence 156, App
	28	246.5	7.0	2680	10	US-09-978-608A-156	Sequence 156, App
	29	246.5	7.0	2680	10	US-09-978-585A-156	Sequence 156, App
	30	246.5	7.0	2680	10	US-09-978-191A-156	Sequence 156, App
	31	246.5	7.0	2680	10	US-09-978-403A-156	Sequence 156, App
	32	246.5	7.0	2680	10	US-09-978-564A-156	Sequence 156, App
	33	246.5	7.0	2680	10	US-09-999-833A-156	Sequence 156, App
	34	246.5	7.0	2680	10	US-09-981-915A-156	Sequence 156, App
	35	246.5	7.0	2680	10	US-09-978-824-156	Sequence 156, App

OM protein - nucleic search, using frame plus p2n model Run on: May 2, 2005, 11:27:24; Search time 5595 Seconds (without alignments) 4496.958 Million cell updates/sec Title: US-09-937-779-2 Perfect score: 3546 Sequence: 1 MGTLFRRNVQNQKSDSDENN......VEPRQLSNLSQWVSQMPFPM 661 Scoring table: BLOSUM62 Xgapop 10.0 , Xgapext 0.5 Ygapop 10.0 , Ygapext 0.5 Fgapop 6.0 , Fgapext 7.0 6.0 , Delext 7.0 Delop 34239544 seqs, 19032134700 residues Searched: Total number of hits satisfying chosen parameters: 68479088 Minimum DB seq length: 0 Maximum DB seq length: 2000000000 Post-processing: Minimum Match 0% Maximum Match 100% Listing first 45 summaries Command line parameters: -MODEL=frame+ p2n.model -DEV=xlh Q=/cgn2_1/USPTO_spool/US09937779/runat 02052005 122715 18269/app query.fasta 1.8 -DB=EST -QFMT=fastap -SUFFIX=rst -MINMATCH=0.1 -LOOPCL=0 -LOOPEXT=0 -UNITS=bits -START=1 -END=-1 -MATRIX=blosum62 -TRANS=human40.cdi -LIST=45 -DOCALIGN=200 -THR_SCORE=pct -THR_MAX=100 -THR_MIN=0 -ALIGN=15 -MODE=LOCAL -OUTFMT=pto -NORM=ext -HEAPSIZE=500 -MINLEN=0 -MAXLEN=2000000000 $-{\tt USER} = {\tt US09937779_ @CGN_1_1_3437_ @runat_02052005_122715_18269} - {\tt NCPU=6} - {\tt ICPU=3} + {\tt USER} = {\tt US09937779_ @CGN_1_1_3437_ @runat_02052005_122715_18269} - {\tt NCPU=6} - {\tt ICPU=3} + {\tt USCR} = {\tt US09937779_ @CGN_1_1_3437_ @runat_02052005_122715_18269} - {\tt NCPU=6} - {\tt ICPU=3} + {\tt USCR} = {\tt US09937779_ @CGN_1_1_3437_ @runat_02052005_122715_18269} - {\tt NCPU=6} - {\tt ICPU=3} + {\tt USCR} = {\tt US09937779_ @CGN_1_1_3437_ @runat_02052005_122715_18269} - {\tt NCPU=6} - {\tt ICPU=3} + {\tt USCR} = {\tt US09937779_ @CGN_1_1_3437_ @runat_02052005_122715_18269} - {\tt UCPU=6} + {\tt UCP$ -NO MMAP -LARGEQUERY -NEG SCORES=0 -WAIT -DSPBLOCK=100 -LONGLOG -DEV_TIMEOUT=120 -WARN_TIMEOUT=30 -THREADS=1 -XGAPOP=10 -XGAPEXT=0.5 -FGAPOP=6 -FGAPEXT=7 -YGAPOP=10 -YGAPEXT=0.5 -DELOP=6 -DELEXT=7 Database : EST:* 1: gb est1:* 2: gb_est2:* 3: gb htc:* 4: gb_est3:* 5: gb_est4:* 6: gb est5:* 7: gb_est6:*

> 8: gb_gss1:* 9: gb_gss2:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

			0			SUMMA	RIES		
_			· 8						
Res		_	Query						
į	No.	Score	Match	Length	DB	ID		Descript	ion
c	1	1274	35.9	948	9	CNS06Y94	puol	71420734	T3 end of
C	2	1188.5	33.5	832	9	CNS060AQ			T7 end of
	3	1020	28.8	978	9	CNS06TH7	1		T7 end of
	4	964	27.2	609	8	BZ297416			CG3420.f1
	5	805	22.7	783	7	CO026714	{		EST805098
_	6		22.7	882	9				
С	7	804.5 732	20.6	899	9 7	CNS07EA9 CO003381			T7 end of
•	8	687	19.4	474	8	AZ928292			EST791716
С	9	685	19.4	400	8				479.dif06
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	13	599.5	16.9	970	7	CO032392			EST810776
	14	586	16.5	512	8	BZ301369	.1/		KD1229.q1
_	15	544.5	15.4	844	6	CB631429	W		OSIIEb08P
С	16	525.5	14.8	550	7	CF884811			tric042xm
	17	513	14.5	854	4	BG645669			EST507288
	18	486	13.7	821	4	BM780050			EST590626
	19	482.5	13.6	724	5	BQ803421			WHE2837_D
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	21	474.5	13.4	719	5	BQ865802			QGC5021.y
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	23	450.5	12.7	867	7	CF821111			EST698493
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	30	426.5	12.0	628	4	BJ478864			BJ478864
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	33	416.5	11.7	595	7	CO135652			EST830323
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	35	416	11.7	902	7	CF704247			CCAC369TR
С	36	414.5	11.7	835	7	CF705715			CCABG95TF
С	37	414.5	11.7	836	7	CF697237			CCAB181TF
С	38	414.5	11.7	858	7	CF705450	,		CCAAF15TF
	39	412.5	11.6	,575	1	AV938810			AV938810
	40	408.5	11.5	610	4	BJ472152			BJ472152
С	41	408.5	11.5	715	7	CF706790			CCAHD40TF
	42	406.5	11.5	596	7	CN846346			PG07016B0
	43	401.5	11.3	731	1	AJ639263			AJ639263
	44	398	11.2	727	6	CD825529			BN25.061A
	45	397.5	11.2	605	2	BE450991		BE450991	EST401878